

Service and Repair Instructions



VENTImotion

BiLevel-ST Home Ventilation Unit VVM 24800

VENTIlogic

BiLevel-ST Home Ventilation Unit WM 27000



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The objective of this service and repair guide is to familiarize you, an **expert, trained specialist,** with the VENTI*motion* and VENTI*logic* in terms of function, technology, servicing and repair. This will enable you to train your customers properly, eliminate faults yourself, perform the function checks specified by the operating instructions and carry out any repairs in accordance with this service and repair guide.

In the event of a claim under warranty, return the devices to Weinmann.

To allow warranty or goodwill applications to be processed, please also send the final customer's proof of purchase (invoice).

Repairs may be performed only by Weinmann or by expert, trained specialists.

You are responsible for repairs carried out yourself and for their warranty!

Use only original Weinmann spare parts for repair.

Remember:

your customer trusts you and is relying on your ability to do the job, just as you rely on Weinmann.

Note

The following information can be found in the operating instructions for the devices:

- safety rules
- setting up device
- operation
- cleaning and disinfecting in use
- warranty







1 Bacteria filter

To protect the device from contamination, especially if the device is being used by several patients.

2 Power supply cable

For connecting the therapy device to the mains power supply.

3 Handle

For transporting the device.

4 Serial interfaces

For connecting to devices, for display, evaluation.

5 Control panel and displays

For controlling and monitoring the therapy device and the connected accessories.

6 Connection for humidifier

For connecting the VENTI*click* humidifier available as an accessory.

7 Device outlet

Respiratory air flows out from here to the patient via the hose system and nasal mask.

8 Sealing plugs (2x)

For sealing the pressure measurement hose during cleaning.

9 Drying adapter

Required to dry the hose system with the therapy device and for function check.

10 Adapter

For connecting the hose system to the device outlet.

11 Pressure measurement hose

For measuring the pressure prevailing in the mask.

12 Hose system

The air flows to the mask through the hose system. The hose system consists of creased hose, pressure measurement hose and adapter.

13 Exhalation system

Carbondioxide-enriched expired air escapes here during therapy.

14 Mask

Respiratory air at the necessary therapy pressure is administered to the patient via the mask.

15 Headgear

For correct and secure positioning of the mask.

16 Filter compartment cover, air inlet

For covering and securely positioning the coarse and fine dust filter.

17 Supply connection

This is where the power supply cable is attached to the device.

18 Cable securing clip

Prevents the device being disconnected from the power supply inadvertently.

19 Connection for rechargeable battery VENTIpower

For connecting the VENTIpower mobile power supply available as an accessory.

20 Connection for O2supply valve VENTI-O2

For connecting the VENTI-O₂ oxygen supply valve available as an accessory.

21 Alarm acknowledgement key with LED

The alarm acknowledgement key is for temporarily muting alarms. The LED provides a visual display of alarms.

22 On/off key

For switching the therapy device on and off.

23 Dial

Central control of the therapy device, for navigating in the menu.

24 Operating keys

For rapid setting by a physician, disabled in patient mode.

25 Menu key

For switching to and fro between the standard display and the menu.

26 Humidifier key with LED

For switching the humidifier on and off or for setting humidifier stage. Six levels are available. The LED indicates whether the humidifier is activated.

27 Softstart key/manual start of analysis (VENTIlogic only)

For activating Softstart and for setting Softstart time up to the maximum Softstart time set in the menu.

In TA mode (VENTI*logic* only), this key is used to start an analysis phase manually.

28 VENTIpower

Available as an accessory to provide a mobile power source for the therapy device.

29 Rating plate

Provides information about the device, such as serial number and year of manufacture.

30 VENTIclick

Available as an accessory for humidifying and heating respiratory air.

31 Carrying bag

For transporting the therapy device.

32 VENTI-O₂

Available as an accessory, for introducing oxygen to the mask.

71 Service label

Indicates when maintenance is next due.

72 Safety test label (in Germany only)

Indicates when the next safety check in accordance with §6 of the German law relating to owners of medical devices [Medizinprodukte-Betreiberverordnung] is due.

Standard display during therapy



1 Status line

This is where information about the status of the device is displayed, such as filter change or maintenance due.

2 Padlock symbol

Indicates whether physician functions are enabled (padlock open) or disabled (padlock closed). The padlock is not displayed in patient mode.

3 Respiratory phase switch indicator

Indicates whether the current respiratory phase switch is spontaneous or mandatory (spontaneous: S, mandatory: T); the indicator switches from left (inspiration) to right (exhalation) depending on respiratory phase; spontaneous exspiration is shown here.

4 Access to menu

The key next to this menu item is used to switch to and fro between the menu and the standard display.

5 Bar chart for pressure indicator

For graphical display of pressure.

6 Ventilation parameters

The relevant current ventilation parameters are displayed depending on the active mode.

7 Active ventilation mode

The active ventilation mode is displayed at this point in the status line.

Symbols used in the display

Symbol	Significance					
Status line:						
-110	Softstart active, remaining time faded in					
\otimes	Filter change required					
4	Maintenance required					
潋	Acoustic signal for the IPAPmin and $V_{T}min$ alarms mute					
λ	Alarm for IPAPmin and V _T min alarms deactivated					
ъ	Physician functions enabled					
â	Physician functions disabled					
×	Fan off (with VENTI <i>motion</i> , only available from software version 6.0)					
Main window						
\bigtriangleup	Low-priority alarm triggered					
$\bigtriangleup \bigtriangleup$	Medium-priority alarm triggered					

Abbreviations used in the display

Symbol	Significance					
Status line:						
TA	TA mode active (VENTI <i>logic</i> only)					
S	S mode active					
ST	ST mode active					
Т	T mode active					
SX	SX mode active					
SXX	SXX mode active					
CPAP	CPAP mode active					
+V	Volume compensation activated (follows mode: e.g. $\mathbf{SXX}^{+\vee}$)					
A ^A	Device in TA mode, automatic analysis phase in progress (VENTI <i>logic</i> only)					
A ^M	Device in TA mode, manual analysis phase in progress (VENTI <i>logic</i> only)					
Main window (m	ionitor)					
IPAP	Inspiration pressure					
EPAP	Expiration pressure					
hPa	Pressure shown in hectopascal; 1.01973 hPa correspond to 1 cm $\rm H_{2}O$					

Symbol	Significance
f	Respiratory frequency
S	Spontaneously-triggered respiratory phase switch
Т	Compulsorily-triggered respiratory phase switch
Main window ("S	tatistics" menu item):
Δр	Maximum rise in pressure due to volume compensation
v	Volume
MV	Per minute volume
Leak	System leak (mask, exspiration system, hose system, device)
f	Respiratory frequency
Ti/T	Proportion of inspiration time in a respiratory cycle
Sins	Proportion of spontaneous inspiration (only when mean values displayed)
Sexs	Proportion of spontaneous exspiration (only when mean values displayed)



Menu structure in Physician mode

2. Description of device

2.1 Intended use for VENTImotion/VENTIlogic

VENTI*motion* and VENTI*logic* are home ventilation devices for the non-invasive, non-life support ventilation of adult patients with respiratory insufficiency in whom there is evidence of autonomous respiratory drive. This corresponds to the following clinical pictures:

- restrictive and obstructive ventilation disorders like paresis of the diaphragm, OSAS, COPD
- disorders of the respiratory mechanism like scoliosis, deformity of the thorax
- neurological, muscular and neuromuscular disorders
- central respiratory regulation disorders

VENTI*motion* and VENTI*logic* are **not suitable for life-support use.**

Use the device only for the purpose described here.

2.2 Functional description

Provision of therapy pressure

An electronically-controlled fan draws in ambient air through a filter and delivers it to the device outlet. From here, the air flows through the hose system and the mask to the patient.

Therapy modes

The device can be operated in the following therapy modes: CPAP, S, ST, T, SX, SXX, TA (TA with VENTI*logic* only).

In time-controlled mode T and in assisted-controlled mode ST, your doctor can set respiratory frequency in the range from 6 to 45 breaths per minute and inspiration time in the range from 20 % to 67 % of the respiratory period.

In assisted modes S, SX and SXX and in assistedcontrolled mode ST, your doctor can select one of 6 trigger levels for both inspiration and exhalation. Your doctor can switch off the trigger for exhalation. Exhalation is then time-controlled.

Other functions

The Softstart function makes it easier to fall asleep. Your doctor sets initial pressures for inspiration and exhalation which continuously rise to the therapy pressures during the Softstart phase. This function can be disabled by the doctor.

The therapy device has an auto switch-on system. If this is activated, the device can be switched on Sensors detect the pressure in the mask and in the hose system and also the respiratory phase switch (trigger point). The fan accordingly provides the IPAP and EPAP pressures set by the doctor.

In the adaptive, controlled TA mode, the device automatically adapts to your personal breathing rhythm and supplies the therapy pressure in exactly the same rhythm.

If no breath is taken into the device in S mode, pressure is switched at a minimum frequency of 6 breaths per minute.

You can activate volume compensation. Minimum volume and maximum pressure rise are set for this purpose. If the minimum volume is undershot, the device automatically increases pressure continuously until the set maximum pressure (therapy pressure + max. pressure rise) is reached.

by a breath being taken into the mask. The device is still switched off using the on/off key (\mathbf{D}) .

The display shows therapy mode and, depending on mode, the values currently being applied for CPAP/IPAP and EPAP and for respiratory frequency (f). Spontaneous or mechanical respiratory phase switches are also displayed and the pressure change shown in the form of a graph.

3.1 Cleaning and disinfecting in use

Caution!

This item is described in section "5. Hygienic treatment" of the therapy device operating instructions.

There follows a description of the hygiene treatment of the device in the event of repair and in the event of a patient change.

3.2 Cleaning and disinfecting on repair

A dealer should perform the following during a repair!

Caution!

It is essential to follow the instructions issued by the manufacturer of the disinfectant (9.3, page 54). It is recommended that you use suitable gloves for disinfecting (e.g. household rubber gloves or disposable gloves).

- Wipe down outer housing and power supply cable with TERRALIN.
- Clean hose, headgear and nasal mask in accordance with the operating instructions or replace with new parts (depending on condition).
- Open device as per item 7.2.
- Replace filters (coarse dust and fine filter).
- Clean out the inside of device housing and filter housing using a vacuum cleaner, clean extremely soiled areas.
- Close device as per item 7.3.

3.3 Cleaning and disinfecting for a patient change

If the device is to be hygiene-treated for another patient, perform the following steps.

Caution!

It is essential to follow the instructions issued by the manufacturer of the disinfectant (9.3, page 54). It is recommended that you use suitable gloves for disinfecting (e.g. household rubber gloves or disposable gloves).

- Wipe down outer housing and power supply cable with TERRALIN. Dispose of hose and mask system, headgear, exhalation system and carrying bag WM 24888 and replace with new parts.
- Open device as per section 7.2.
- Clean out the inside of the device housing, box and filter holder using a vacuum cleaner, clean extremely soiled areas.

- Replace coarse and fine dust filters **39** + **40**.
- Open box **58** as per section 7.7.
- Spray-disinfect box, lid, fan, fan flap, motor bearing, fan cable, filter holder 37 and filter compartment cover 16 twice with MIKROZID LIQUID, in each case waiting the prescribed time for the product to take effect. In addition, at the start of the time for the product to take effect, wipe down accessible areas with a cloth wetted in MIKROZID LIQUID.
- Replace the following parts with new parts:
 - hose system WM 24130
 - filters WM 24870 and WM 24880
 - pressure and flow measurement hoses
 WM 24324, WM 24835
 - hose, pressure side WM 24852

- device outlet, complete, WM 24025
- labyrinths WM 24897
- decoupling tube WM 24028
- motor frame WM 24898
- cushioning insert, box WM 24874
- fan box gasket WM 24868
- filter holder gasket WM 24151
- hose, intake side WM 24809
- foam for expansion space WM 24171
- foam for filter holder WM 24155
- countersunk screw ISO 7045-M4x6-V2A-H with Tuflok
- Seal box **58** again as per section 7.7.
- Alternatively, the box 58 can be replaced by a reconditioned box as described in section "7.6 Replace box and filter holder" on page 35.

- Insert device outlet WM 24025
- Replacing the WM 24837 control board Note: This is necessary because the flow sensor located on the control board cannot be sterilised.
- Close device as per item 7.3, disinfect outer housing and power supply cable with TERRALIN wipes.
- Reset maintenance symbol to set the maintenance indicator to "O" (see "5.4 Reset maintenance symbol" on page 26).
- Test the device.
- Delete patient data stored in the device as described in the hospital manual for the VENTImotion.

3.4 Clean and disinfect humidifier during use

This item is described in section "4. Hygiene treatment" in the VENTIclick operating instructions.

3.5 Clean and disinfect humidifier on patient change

If the device is to be hygiene-treated for another patient, perform the following steps.

- For hygiene reasons we recommend replacing plastic parts after a maximum period of use of 2 years. The spare parts list can be found in the operating instructions for the VENTIclick.
- If plastic parts and the heating element are heavily soiled or coated in limescale, offer a new device, otherwise: proceed in accordance with section "4. Hygiene treatment" in the operating instructions for VENTIclick.

3.6 Clean and disinfect the VENTIpower

This item is described in section "4. Hygiene treatment" in the VENTIpower operating instructions.

3.7 Clean and disinfect the VENTI-O2

This item is described in section "4. Hygiene treatment" in the VENTI-O₂ operating instructions.

4.1 General

Important!

The device must be subjected to the following test following every repair, maintenance and hygienic treatment in accordance with test instruction WM 24827 and the test protocol. The test can also be used to assist during troubleshooting.

For the therapy device, enter the operating hours and all parameters in your service protocol (see page 59). If you find faults or deviations from target values during the test, you must not use the therapy device again until the faults are eliminated.

The possible causes of the faults and how you eliminate malfunctions can be found in section "6. Malfunctions and rectification" on page 28.

4.2 Test equipment

- Flow measurement device
- Test set WM 23465 incl. 23635 pressure measurement adapter
- PC and PC software VENTIsupport
- Converter box WM 93316
- Connecting cables WM 93312 and WM 96313
- 115 V power source, e.g. travel adapter 230 V/115 V, 200 W
- Test protocol (see "12.2 Test protocol" on page 60)

4.3 Preparation for testing

Check power supply cable

1. Check the power supply cable 2.

Ensure that

- the insulation is OK,
- the cable is undamaged
- and there are no loose contacts.
- 2. Replace the power supply cable **2** if necessary.

Check housing

Check the housing for general condition.

If the housing is damaged or defective, replace the relevant side of the device (see "7.13 Replace bottom part of housing" on page 46/"7.14 Replace top part of housing" on page 47).

Connection to VENTIsupport

- 1. Plug the plug of the power supply cable into a socket.
- 2. Have the test protocol to hand.
- 3. Connect the serial interface of the therapy device to the PC via the converter box.
- 4. Start the VENTI*support* program.
- 5. Click on the **VENTIadjust** item in the menu bar.

VENTIadjust								
ta Therapy param	eter setting	Tec	hnical data	setting Alarm settin	gs			
Therapy parameters								
Therapy device	VE	NTII	logic		AF:	12	/min	
Therapy device:	Au	s			Ti/T:	33	%	
Therapy mode:	ST	mod	de		Insp. trigger:	3		
Actual pressure:	0		hPa		Exp. trigger:	3		
EPAP pressure:	4		hPa		Pressure rise insp.:	1		
IPAP pressure:	6		hPa		Pressure rise exp.:	1		
Soft start pressure E	PAP: -		hPa		Volume compensation	Off		
Soft start pressure IF	AP: -		hPa		VT:	300	ml	
Softstart:	-				Delta P:	5	hPa	
			min		Autostart:	Off		
Softstart time:	-							
Softstart time: Soft start time Tmax:	-		min		Humidifier:	Off		
Softstart time: Soft start time Tmax: Soft start disabled:	- - 0N	1	min		Humidifier: Humidifier stage:	Off 2		
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since:	- - 0N		min	07.06.2006	Humidifier: Humidifier stage: Therapy duration/day::	Off 2		0 h
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since: Therapy duration:		 	min	07.06.2006 0 h	Humidifier: Humidifier stage: Therapy duration/day:: Therapies/day:	Off 2		0 h 0
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since: Therapy duration: Therapy days	0N	 	min	07.06.2006 0 h 0	Humidifier: Humidifier stage: Therapy duration/day:: Therapies/day:	Off 2		0 h 0
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since: Therapy duration: Therapy days Service	- - ON			07.06.2006 0 h 0	Humidifier: Humidifier stage: Therapy duration/day: Therapies/day:	Off 2		0 h 0
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since: Therapy duration: Therapy days Service Date/Time:		14:28	3.07	07.06.2006 0 h 0	Humidifier: Humidifier stage: Therapy duration/day:: Therapies/day:	Off 2		0 h 0
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since: Therapy duration: Therapy days Service Date/Time: Fine filter:		14:28	min	07.06.2006 0 h 0	Humidifier: Humidifier stage: Therapy duration/day:: Therapies/day:	Off 2		0 h 0
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since: Therapy duration: Therapy days Service Date/Time: Fine filter: Operating hours:		14:28	3:07	07.06.2006 0 h 0	Humidifier: Humidifier stage: Therapy duration/day: Therapies/day:	Off 2		0 h 0
Softstart time: Soft start time Tmax: Soft start disabled: Device usage Patient since: Therapy duration: Therapy duration: Therapy days Service Date/Time: Fine filter: Operating hours: SW version:		14:28	3:07	07.06.2006 0 h 0	Humidifier: Humidifier stage: Therapy duration/day: Therapies/day:	Off 2		0 h 0

6. Note the set therapy parameters.

4.4 Enter device data

• Enter the type, number and date of manufacture of the device in the test protocol.

4.5 Test temperature calibration and calibration status

Note

This test is not needed, as it can only be performed on a Weinmann test bench.

- 1. Plug the red drying adapter into the device outlet of the VENTI*motion*.
- 2. Turn on the device.

Requirement: if battery voltage is sufficient, **no** maintenance symbol should appear in the display.

- 3. Replace
- 4. Turn off the device.

Note

If battery voltage is inadequate, replace the battery on the control board (see "7.9 Replace battery on control board" on page 42).

4.7 Test interface with the VENTIclick

• Check the function of the humidifier (see " Check function of humidifier" on page 21).

4.8 Check target pressure values 5, 20 and 35 hPa

- 1. Connect the hose system and the mask to the therapy device.
- 2. Pull a plug off the mask.

- 3. Connect a pressure meter to the opened connector.
- 4. Turn on the device.
- Prepare the therapy device for testing by making the following settings in the "Physician" menu:
 - T mode
 - Frequency f = 6/min
 - Ti:T = 50 %
 - EPAP 5 hPA
 - IPAP 35 hPA



- 6. Read off the pressure displayed by the measurement device and enter it on the protocol in the "Measuring result" column.
- 7. Read off the actual pressure displayed by the therapy device and enter it on the protocol in the "Mask" column.
- 8. Change the setting from EPAP 5 hPA to EPAP 20 hPA.
- Read off the pressure displayed by the measurement device and enter it on the protocol in the "Measuring result" column.
- 10. Read off the actual pressure displayed by the therapy device and enter it on the protocol in the "Mask" column.
- 11. Turn off the device.

Requirement: the values must be within the tolerances quoted in the protocol.

Note

If the values are not within the quoted tolerances, replace the control board (see "7.8 Replace control board/display" on page 39).

4.9 Test flow measurement

- Connect the device output of the therapy device to the flow measurement device using the short hose.
- 2. Turn on the device.
- 3. Prepare the VENTI*motion* for testing by making the following settings in the "Physician" menu:
 - T mode
 - frequency f = 6/min

4. Set the two pressures so that the device displays 50 and 100 l/min in the flow window. For example: IPAP=10 hPa and EPAP=5 hPa Call up the flow window by pressing the rotary knob in the "Physician" menu during operation and selecting "Flow curve" in the Display menu.

Correct the entry for pressures if necessary.

- Enter the values displayed by the flow measurement device in the protocol.
 Requirement: the values must be within the tolerances quoted in the protocol.
- 6. Turn off the device.

Note

If the values are outside the tolerances, replace the control board (see "7.8 Replace control board/display" on page 39).



4.10 Test leaktightness

- 1. Plug the test adapter onto the device outlet.
- 2. Connect the side connection to the pressure measurement device.
- Turn on the device. Select CPAP mode in the "Physician" menu and set a pressure of 20 hPa.
- 4. Read off the pressure displayed by the pressure measurement device and enter it in the protocol.

Requirement: the values must be within the tolerances quoted in the protocol.

5. Turn off the device.

Note

If the value is outside tolerance, dismantle the device (see "7.2 Open device" on page 31) and check for leaks.

4.11 Test clock function

- 1. Connect the serial interface of the therapy device to the PC via the converter box.
- 2. Start the VENTI*support* program.
- 3. Click on the **VENTIadjust** item in the menu bar.
- 4. Select the **Set technical data** tab.
- Compare "Device time" and "PC time".
 Requirement: the values must change to the same extent.





Device time Date: 12.09.2006 Time: 10:08:07 PC time Date: 12.09.2006 Time: 10:08:59	~	aces name								
Date: 12.09.2006 Time: 10:08:07 PC time Date: 12.09.2006 Time: 10:08:59	1	Device time								
PC time Date: 12.09.2006 Time: 10:08:59		Date:	12.09.2006	Time:	10:08:07					
Date: 12.09.2006 Time: 10:08:59		– PC time	.							
		Date:	, 12.09.2006	Time:	10:08:59					

4.12 Test the interface with the VENTI- O_2

- 1. Fit a VENTI- O_2 value to the therapy device so that it is ready to work. The indicator on top of the value housing does not light up.
- 2. Close the opening of the nasal mask, using your thumb or hand, for example.
- Switch on the therapy device first, then your O₂ supply. After the therapy device has been switched on, the valve opens with a soft "click" indicated by the green indicator.
- 4. You can now set the test flow rate on the flow display of your oxygen system. If this is not possible, first check the function of your oxygen system (is the cylinder empty, for example, or are hoses kinked?).
- 5. Switch the device off again. The valve audibly switches to "vent", the indicator goes out.

Note

If the VENTI- O_2 value does not react as described, perform the following measures in sequence and test again in each case.

- Use a different valve.
- Replace the connecting cable of the power board/mains supply unit (see "7.5 Replace power board" on page 34).
- Replace the power board (see "7.5 Replace power board" on page 34).

4.13 Test interface with the VENTIpower

Note

Before this test is carried out, the therapy device must have been connected to the mains power supply for at least 5 minutes.

- Fit the VENTIpower to the therapy device so that it is ready to work therapy device. Connect the therapy device to the mains power supply.
- Switch on the VENTIpower and then the therapy device. VENTIpower is working correctly if its displays go out, in other words, if VENTIpower switches to standby mode.
- Now disconnect the plug of the therapy device, the device switches off. The power failure alarm sounds.



VENTI*power* is working correctly if its displays comes on after approx. 4 seconds and the therapy device starts working again.

- Now restore the mains power supply to the therapy device. VENTIpower is working correctly if its displays go out, in other words, if VENTIpower switches to standby mode.
- 5. Switch off both devices.

4.14 Test noise

Note

This test is not needed, as it can only be performed on a Weinmann test bench.

4.15 Test all keys, the rotary knob and 115 V mode

- 1. Plug the red drying adapter into the device outlet.
- 2. Connect the device to a 115 V power source and switch on the device.

Requirement: the device must start working.

- 3. Press all the keys and the rotary knob to test their function.
- 4. Turn off the device.

Note

If the keys do not work properly, perform the following measures in sequence and test again in each case.

- Check fascia film and replace if necessary (see "7.12 Replace fascia film" on page 45).
- Replace the control board (see "7.8 Replace control board/display" on page 39).

Note

If the rotary knob does not work properly, perform the following measures in sequence and test again in each case:

- Change the encoder (see "7.11 Replace encoder (dial)" on page 44).
- Replace the control board (see "7.8 Replace control board/display" on page 39).

Note

if 115 V mode does not work properly, replace the mains supply unit (see "7.4 Replace mains supply unit" on page 33).

4.16 Test display incl. backlighting and contrast, the LEDs and the alarm

Note

Before this test is carried out, the VENTI*motion* must have been connected to the mains power supply for at least 5 minutes.

- Switch on the device. A beep sounds. Both LEDs come on briefly.
- 2. Check whether the display is easy to read with backlighting and that contrast is adequate.

Set contrast

- 1. Disconnect the power supply plug.
- 2. Press the humidifier key <u>≤</u> and the Softstart key **→** simultaneously.
- 3. Connect the power supply plug and the socket again.
- 4. Turn the rotary knob to select contrast. Activate contrast by pressing the rotary knob.

Note

If no beep is heard when you switch on, perform the following measures in sequence and test again in each case.

- Test the buzzer and replace if necessary (see "7.13 Replace bottom part of housing" on page 46).
- Replace the control board (see "7.8 Replace control board/display" on page 39).

Note

If the contrast cannot be set, perform the following measures in sequence and test again in each case.

- Check the display cable.
- Replace the display (see "7.8 Replace control board/display" on page 39).
- Replace the control board (see "7.8 Replace control board/display" on page 39).

4.17 Test equipment and accessories (system components)

- Humidifier VENTIclick present
- Hose system present
- Oxygen supply valve VENTI-O₂ present
- Rechargeable battery VENTIpower present
- Medical devices manual present (Germany only)
- Operating instructions present

Check function of humidifier

Important!

It is essential to note the heating level set before carrying out this test.

Caution

Perform this test even if the patient has not used or will not be using the humidifier.

- Subject the plastic housing to a visual check: in the event of cracks/damage and severe soiling, the plastic parts or gaskets should be replaced.
- 2. Fill the humidifier with water up to the mark.
- 3. Check whether the humidifier is leaktight.
- 4. Pour out the water.
- 5. Now pour in 50 ml of water.
- 6. Click the humidifier onto the therapy device.
- 7. Plug the red drying adapter (included in the scope of supply of the therapy device) into the outlet connector of the humidifier.
- 8. Plug the hose system onto the drying adapter.
- 9. Turn on the therapy device.
- 10. Turn on the humidifier by pressing the humidifier key () on the therapy device.
- 11. Set heating level 6 on the therapy device.
- 12. Check whether the humidifier is heating up.
- 13. Take off the hose system by pushing the locking button of the adapter.
- 14. Pull the red drying adapter out of the humidifier by twisting it slightly.
- 15. Set the heating level back to the value you noted.

Note

If the humidifier has not heated up, perform the following measures in sequence and test again in each case.

• Use a different humidifier. Is the heating element OK?



Replace the humidifier socket and cable (see "7.13 Replace bottom part of housing" on page 46).

• Replace the power board (see "7.5 Replace power board" on page 34).

Function check VENTI-O2

- Fit the VENTI-O₂ to the therapy device so that it is ready to work. The status display on top of the valve housing does not light up.
- 2. Attach the test hose supplied (length approx. 48 cm) to the O_2 outlet (\square).
- First switch on your therapy device and then your O₂ supply. After the therapy device has been switched on, the valve opens with a soft "click". The green status display lights up.
- 4. Close the opening of the mask, using a thumb or hand, for example.
- 5. Set the prescribed flow rate at the flow setting of your oxygen system. If this is not possible, first check the function of your oxygen system (is the cylinder empty, for example, or are hoses kinked?).
- 6. Hold the free end of the test hose in a glass half-filled with water. VENTI- O_2 is working correctly if bubbles escape from the end of the hose.
- Switch the therapy device off again. VENTI-O₂ is working correctly if the valve audibly switches to "vent", the status display goes out and no more bubbles escape from the end of the test hose.
- 8. Shut off the oxygen supply of your oxygen system.

4.18 Once tests are complete

• Reset the therapy values you noted during preparations.



5. Maintenance

5.1 Intervals

Both filters **39** and **40** must be regularly checked for soiling.

- Coarse dust filter **39** needs to be changed every 6 months.
- Fine filter 40 should be changed after no more than 1000 operating hours (filter change symbol appears in the display).

For reasons of hygiene, we recommend replacing the following parts at the intervals stated:

- pressure measurement hose 11 every 6 months or sooner if soiled.
- complete mask system every 6 to 12 months depending on soiling
- exhalation system in accordance with the relevant operating instructions

See section entitled "Cleaning" in the relevant operating instructions.

Servicing should be carried out at the following intervals as a preventive measure:

- after every 5,000 operating hours (maintenance symbol appears in the display and should be reset after every service),
- 2. after no more than 2 years (see service label on rear of device).
- 3. after 10,000 hours or 4 years, clean or replace all parts in the air flow.
- Replace service label **71** with one with the new data (see "8. Spare parts" on page 48). Cut out the month at a sharp angle using a punch or nail scissors. Affix the new service label to the left-hand side next to the filter flap.



In **Germany**, a safety control as laid down in §6 of the German law relating to owners of medical devices [Medizinprodukte-Betreiberverordnung] must also be carried out every 2 years as specified.

5.2 Filter change

Use only original filters made by Weinmann. If third-party filters are used, this will invalidate the warranty and may cause restrictions in terms of function and biocompatibility.

Coarse dust filter

- If the VENTIclick humidifier is connected, first disconnect it from the device. This will prevent water running into the device when the filter is changed. Please also see the operating instructions for the VENTIclick.
- 2. Press on the latch of the filter compartment cover and lift it off.
- 3. Take coarse dust filter **39** out of the filter compartment cover and dispose of it with ordinary domestic waste.
- 4. Insert clean coarse dust filter **39** in the filter compartment cover.
- 5. Insert the filter compartment cover into the opening in the housing, bottom edge first. Then push the filter compartment cover into the housing until the latch engages.

Fine filter

The fine filter needs changing when it has turned dark in colour, but in any event, after no more than 1000 operating hours. In the latter case, when VENTI*motion* is switched on, the message "Change filter!" appears in the display.

Acknowledge the message by pressing the alarm acknowledgement key 🖄. The filter change symbol 🞆 then appears continuously in the status line. To change the fine filter, proceed as follows:

- 1. Press on the latch of the filter compartment cover and lift it off.
- 2. Take out fine filter **40** and dispose of it with ordinary domestic waste.
- 3. Insert a new fine filter **40** WM 15026.
- Insert the filter compartment cover into the opening in the housing, bottom edge first. Then push the filter compartment cover into the housing until the latch engages.
- 5. Reset the filter change indicator (see "Reset filter change indicator" on page 25).









Reset filter change indicator

- To reset the filter change indicator, press menu key in with the device switched on and use the dial to select the item Filter change in the Patient menu or in Physician mode, select the menu item Device configuration and then Filter change. Push the dial to call up the "Filter change" menu.
- The question "Reset filter change?" appears. Use the dial to select YES and confirm your selection by pressing the dial.

If you wish to cancel the process, use the dial to select **NO** and press the dial. The process is aborted.

Once you have selected and confirmed **YES** with the dial, the message "Filter change reset!" appears for about 3 seconds.

Bacteria filter

If bacteria filter WM 24148 is used, change the particle filter in the bacteria filter in accordance with the relevant operating instructions.

5.3 Device cleaning

The parts in the air flow should all be cleaned and disinfected every 10,000 hours or every 4 years. This should be performed in accordance with section "3.3 Cleaning and disinfecting for a patient change" on page 11.

After every maintenance operation, perform a test according to section "4. Test the device" on page 13.



After every service/repair performed, the maintenance indicator must be set to "O", or the maintenance symbol which has appeared in the display deleted. In addition, a new service label (current year +2 years) should be affixed to the rear of the device.

- Enable Physician functions. Hold down the IPAP and EPAP keys for 4 seconds to do so. The message "This is a physician function!" appears in the display. The display then returns to Monitor. The status line of the display shows the symbol a. The Physician functions are now enabled.
- 2. In the **menu**, select the sub-menu **Device configuration**.
- Use the dial to select the menu item Maintenance. Confirm your selection by pressing the dial.

The Maintenance sub-menu is displayed.

If you wish to reset the indicator, use the dial to position the selection bar on **YES**. Confirm your selection by pressing the dial.

- A display appears in which you are asked again whether you wish to reset the symbol. If you are sure, select YES here too and confirm your selection by pressing the dial. The message: "Maintenance reset!" appears for approx. 2 seconds.
- To exit the menu, press the menu key (back) until the Monitor is displayed. You can also select back with the dial and then press the dial.



When you have maintained the device, a new maintenance sticker (current year +2 years) must be attached to the rear of the device.

• Replace maintenance sticker **71** with one showing the new data (see "8. Spare parts" on page 48). Cut out the month using punch pliers or the tip of a nail cutter. Affix the new maintenance sticker to the left of the filter flap.



5.6 Disposal



Do not dispose of the device with domestic waste. To dispose of the device properly, please contact a licensed, certified electronic scrap disposal merchant. This address is available from your Environment Officer or from your local authority.

6.1 General faults

Malfunction	Cause	Rectification	
	No power supply.	Check connection of power supply cable in device socket. Check mains voltage with another device, if necessary (for example a lamp), replace power supply cable if necessary.	
No running noise standby	Mains supply unit defective.	Replace mains supply unit (7.4, page 33).	
and operating displays do not light up.	Cable come loose or defective.	Check all cable connections in the device; replace if necessary.	
	Control board defective.	Replace control board (7.8, page 39).	
	Power board defective.	Replace power board (7.5, page 34). Send defective board to the manufacturer for fault analysis.	
	Display defective.	Replace display (7.8, page 39).	
Faulty displays or none at all	Cable come loose or defective.	Check all cable connections in the device; replace if necessary.	
	Control board defective.	Replace control board (7.8, page 39).	
Tolorance of thorapy prossures	Control board defective.	Replace control board (7.8, page 39).	
is > 0.6 hPa after 1 minute.	Sensor hoses defective or soiled.	Check hoses, lay correctly (7.8, page 39).	
In Automatic mode, device	Automatic system not active	Activate auto switch-on/switch-off (see 4.1 of operating instructions).	
can not be switched on by a breath being taken in.	Power board defective.	Replace power board (7.5, page 34).	
	Control board defective.	Replace control board (7.8, page 39).	
Device is running but does	Filter soiled.	Change both filters (5.2, page 24).	
not reach the lower pres-	Nasal mask leaking.	Adjust headgear/headband so that the mask is tight.	
sure limit set.	Leak in device.	Check if all hoses and gaskets are properly located.	
Filter change indicator	Filter soiled.	Clean/change both filters (5.2, page 24).	
Maintenance display	Maintenance interval exceeded.	Perform maintenance.	
active.	Internal battery discharged.		
	No power.	Check that plug contact is properly located.	
Status display of the VENTI-O ₂ lights up intermit- tently or not at all.	Nasal mask leaking.	Adjust headgear/headband so that the mask is tight or use another mask if necessary.	
	Hose leaking.	Check breathing hose and pressure measurement hose.	
	O ₂ supply hoses kinked.	Check all hose connections.	
The prescribed O ₂ flow is not reached.	Too high a resistance in the hose system.	Shorten O_2 supply hoses.	
	Output pressure of oxygen system too low.	If necessary, use another oxygen system, e.g. OXYMAT 3.	
Water in humidifier is not	Humidifier defective.	Test with a different humidifier, if device is defective, return humidifier to the manufacturer.	
heating up.	Power board defective.	Replace power board (7.5, page 34). Send defective board to the manufacturer for fault analysis.	
Power supply cable dam- aged.		Replace power supply cable.	

Device cannot be switched on using the film keypad.	The ribbon cable of the film keypad has slipped out of the connection.	Check whether the ribbon cable is correctly connected to the display board.
	Fascia film defective.	Replace fascia film (7.12, page 45).

6.2 Internal faults

The therapy device performs a self-test of the sensor system when the on/off switch igodot is pressed briefly, checking the function of the alarm. If a fault occurs during the self-test, an error message appears in the main screen (see also "6.2 Internal faults" on page 29).

The therapy device can detect internal faults itself and shows them on the display. The display takes the form either of clear text or of a code.

Fault output in clear text

Display	Cause	Rectification	
"Battery discharged!"	Internal battery of the device is discharged.	Change the battery (7.9, page 42).	
"Clock not set!"	Time not set on device following battery change	Set clock as described in VENTI <i>motion</i> hospital manual.	
Device error; excessive pressure	Pressure at mask or fan has exceeded maximum permitted limit value.	Check ventilation and pressure measurement hose system for blockages/leaks.	
	Pressure measurement hose blocked		
Device error, Upper temperature limit exceeded (Code 64)	The temperature on the power board has reached a critical value, e.g. as a result of opera- tion in the bag or being in direct sunlight.	Leave device to cool down and operate it in the permitted ambi- ent temperature range.	

Coded error messages

For faults which are not displayed with a clear text message, the therapy device issues a fault code.

- A number appears in the first line.The appropriate 16-digit fault code appears in the second line. Each of the 16 digits has the value "O" or "1". The position of the 1 indicates the kind of fault involved.

A description of the cause of the fault and how to rectify it can be found in the table below.

Note: several errors may occur simultaneously. In this case, a "1" will then appear at several points in the second line. The number in the first line is the total of all the numbers for individual faults.



First line: code number	Second line: 1 at position	Cause	Rectification
			Replace battery on control board (see section 7.9 on page 42).
1	1	Clock not responding	If fault occurs again, replace control board (see section 7.8 on page 39).
		battery discharged	Replace fan or fan box. (see section 7.7 on page 37).
			Replace power board (see section 7.5 on page 34).
2	2	EEPROM not responding	Replace control board (see section 7.8 on page 39).
4	3	Transmitting fault at the serial connections.	Replace control board (see section 7.8 on page 39). If this occurs on final test bench, please inform QM.
8	4	Sensor measurement has failed plausibility test.	Replace control board (see section 7.8 on page 39).
		Fan has failed or fan not	Check cable connections between fan, power board and control board.
16	5	starting up. Either there is no 40 V supply or there is	Replace fan or fan box (see section 7.7 on page 37).
		a detect on the tan.	Replace power board (see section 7.5 on page 34).
128	8	Humidifier current is out- side permitted values.	Replace power board (see section 7.5 on page 34).
256	9	EEPROM has failed checksum test.	Replace control board (see section 7.8 on page 39).
512	10	Sensor calibration data in EEPROM invalid.	Replace control board (see section 7.8 on page 39).
1024	11	Pressure sensor measured	Check external and internal pressure meas- urement hoses for blockages.
		values outside tolerance	Replace control board (see section 7.8 on page 39).
2048	12	There are invalid therapy parameter values in the EEPROM.	Replace control board (see section 7.8 on page 39). If this occurs on final test bench, please inform QM.
4096	13	Data inconsistency, possi- bly battery discharged	Replace battery on control board (see section 7.9 on page 42).
8192	14	No communication with EEPROM.	Replace control board (see section 7.8 on page 39).
16384	15	Communication with watchdog failed.	Replace control board (see section 7.8 on page 39).
32768	16	Software fault	Replace control board (see section 7.8 on page 39).

7. Repairguide for VENTImotion/VENTIlogic

7.1 General

Perform repairs on the VENTImotion or VENTIlogic exclusively at an ESD protected work-station!

- Follow the safety rules in the operating instructions for the therapy device.
- Any handling of the device assumes accurate knowledge and observance of the operatinginstructions and the service and repair guide.
- Only perform repairs which are described in this service and repair guide. This is the only way to guarantee that the therapy device continues to function properly.
- Ensure that your hands and your work-station are clean during repair work.

- Perform a function check after every repair (see "4. Test the device" on page 13).
- If you replace components or individual parts, use only original Weinmann parts.
- When ordering the bottom part of the housing **35**, quote the type, year of manufacture and number of the device as well.

Note

The item numbers listed in the text which follows are identical to the item numbers in the spare parts list on page 48 and in the overview on page 4.

7.2 Open device

Tools and equipment required

- ESD protected work-station
- 3 mm Allen key

Caution!

Open the device only with the power supply plug disconnected.

- 1. Place the device on a non-slip surface with the top facing downwards.
- 2. Remove the filter cassette from the top part of the housing.
- 3. Undo and remove the 6 screws 34.



- 4. Open the bottom part of the housing **35** away to the side.
- 5. Undo the connecting cables for the humidifier **42** and the alarm **43**.
- 6. You can now put the bottom part of the housing **35** to one side.



7.3 Close the device

Tools and equipment required

- ESD protected work-station
- 3 mm Allen key
- 1. Hold the bottom part of the housing **35** up to the side of the top part of the housing **36**.
- 2. Plug the connecting cables for the humidifier **42** and the alarm **43** onto the relevant connectors.
- Check that all hoses and cables are plugged on firmly. If necessary, carefully plug these fully onto the connections (hoses) or slots (cables).
- 4. Place the bottom part of the housing **35** on the top part of the housing **36**.

Ensure that no cables or hoses are trapped or bent.

- 5. Now screw the top part of the housing tight using the 6 screws **34**.
- 6. Then turn the device back over.





- ESD protected work-station
- 3 mm Allen key
- 1. Open the device (see "7.2 Open device" on page 31).
- Disconnect the power board/mains supply unit connecting cable 45 from the power board (wide connector).
- 3. Slightly lift mains supply unit **44** and disconnect the mains input connector from the mains supply unit **44**.
- 4. Lift the mains supply unit out of the top part of the housing.
- Disconnect the power board/mains supply unit connecting cable from the mains supply unit and plug into the new mains supply unit.
- 6. Position the new mains supply unit in the top part of the housing. Plug the mains input connector onto the new mains supply unit in the process.
- 7. Plug the power board/mains supply unit connecting cable **45** onto the power board (wide connector).

Caution!

Be absolutely sure that power board/mains supply unit connecting cable 45 is between the head of the screw and the box as well as in the groove of the power board. Ensure that the rechargeable battery cable is in the groove. Otherwise the cable may be trapped and damaged when the device is closed.

- 8. Close the device (see "7.3 Close the device" on page 32).
- 9. Test the device (see "4. Test the device" on page 13).



- ESD protected work-station
- 3 mm Allen key
- 1. Open the device (see "7.2 Open device" on page 31).
- Disconnect the power board/mains supply unit connecting cable 45 from the power board 41 (wide connector).
- 3. Disconnect the motor cable.
- Disconnect the O₂ valve/power board connecting cable **70** from the power board.
- 5. Disconnect all connecting cables from the control board.
- 6. Remove the power board from the top part of the housing.
- 7. Take the rechargeable battery connecting cable **48** out of the groove and disconnect it from the power board **41**.
- Disconnect the cables of the old power board and plug them into the appropriate slots of the new power board 41:
 - power board/control board connecting cable 46 (wide connector)
 - rechargeable battery connecting cable 48
 - earth cable 47
- 9. Connect the power board/mains supply unit connecting cable **45** to the power board.
- 10. Position the new power board in the top part of the housing.
- 11. Restore the connections to the control board.
- 12. Connect the O_2 value connecting cable to the new power board.
- 13. Plug the motor cable back on.







14. Insert the rechargeable battery connecting cable into the groove on the power board.

Caution!

Be absolutely sure that the power board/mains supply unit connecting cable 45 is between the head of the screw and the box as well as in the groove of the power board. Ensure that the rechargeable battery cable is in the groove. Otherwise the cable may be trapped and damaged when the device is closed.

- 15. Close the device (see "7.3 Close the device" on page 32).
- 16. Test the device (see "4. Test the device" on page 13).

7.6 Replace box and filter holder

Tools and equipment required

- ESD protected work-station
- 3 mm Allen key
- 1. Open the device (see "7.2 Open device" on page 31).
- 2. Remove the power board (see "7.5 Replace power board" on page 34).
- Remove the mains supply unit (see "7.4 Replace mains supply unit" on page 33).
- 4. Release the pressure-side hose **74** (connection to device outlet) from the box.
- 5. Lift box **58** and filter holder **37** out of the top part of the housing together.
- Disconnect the box and the filter holder from one another by undoing the intake-side hose 73 from both parts.





7. Dispose of the foam **38** from the expansion space and replace box **58**.

or:

Replace the complete filter holder **37**.

- 8. If you have not replaced the filter holder, insert a new piece of foam **38** in the expansion space.
- Connect box and filter holder. To do so, insert the intake-side hose in both parts. The smooth part of the hose must be inserted in the filter holder in each case.
- Insert the box and the filter holder back in the device (steps 4. to 6. in reverse sequence). To facilitate assembly of the intake-side hose 73, apply a little 70 % isopropanol to the circumference.
- 11. Refit the mains supply unit (see "7.4 Replace mains supply unit" on page 33).
- 12. Refit the power board (see "7.5 Replace power board" on page 34)
- 13. Close the device (see "7.3 Close the device" on page 32).
- 14. Test the device (see "4. Test the device" on page 13).



- ESD protected work-station
- 3 mm Allen key
- 1. Open the device (see "7.2 Open device" on page 31).
- 2. Remove the power board (see "7.5 Replace power board" on page 34).
- 3. Remove the mains supply unit (see "7.4 Replace mains supply unit" on page 33).
- 4. Remove box and filter holder (see "7.6 Replace box and filter holder" on page 35).
- 5. Open the box. To do so, lift the lid using a slot screwdriver. Then remove the lid of the box.
- 6. Lift fan **63** up and disconnect decoupling tube **66** from the bore of the motor frame.
- 7. Remove fan 63 from box 58.

If the same patient is going to be using the device, continue the repair from step **15**.

If box **58** is to be treated for a new patient or be subjected to cleaning in the course of the 10,000-hour/4-year service, then proceed as follows.

- 8. Remove motor frame **60** and the labyrinths **59** from the box and dispose of them.
- 9. Remove lid cushioning insert **61** and box cushioning insert **62** from the box and dispose of them.
- 10. Clean out the box using a vacuum cleaner and clean extremely soiled areas.
- 11. Disinfect the box using MIKROZID LIQUID (follow the manufacturer's instructionsfor use).







 Insert a new lid cushioning insert 61 and box cushioning insert 62 as shown, as well as the two new labyrinths 59 and motor frame 60 in box 58.

Take care with the assembly position.

- Insert the labyrinths as shown.
- Insert motor frame 60 so that the cutout for the fan cable is flush with the cutout in the box. Otherwise the fan cable might be damaged when the box is closed.

13. Tip!

Put motor frame **60** and labyrinths, **59** together with box cushioning insert **62**, in box **58**. Clean the fan and then disinfect it using MIKROZID LIQUID (follow the manufacturer's instructions).

- 14. Fit the fan flap with two new screws **68** from the maintenance kit.
- 15. Insert fan **63** in motor frame **60**.
- 16. Apply a little 70 % isopropanol to the circumference of decoupling tube 66 and fit it in the appropriate bore of motor frame 60. The groove of the hose must engage right round the circumference of the bore.
- 17. Put the lid on box 58.

Ensure that the lid is fitted in the correct position. The cable harness of the fan must run through the cutout in the lid and may not be trapped! The cable must protrude between 265 and 270 mm.

- 18. Refit the box and filter holder (see "7.6 Replace box and filter holder" on page 35)
- 19. Refit the power board (see "7.5 Replace power board" on page 34).
- 20. Refit the mains supply unit (see "7.4 Replace mains supply unit" on page 33).
- 21. Close the device (see "7.3 Close the device" on page 32).
- 22. Test the device (see "4. Test the device" on page 13).





Perform repair

Tools and equipment required

- ESD protected work-station
- 3 mm Allen key
- Phillips screwdriver, size 1
- 1. Open the device (see "7.2 Open device" on page 31).
- 2. Remove the power board (see "7.5 Replace power board" on page 34).
- Disconnect the encoder connector 82 and the connectors of interfaces 84 and 85 of the control board.

 Disconnect the ribbon cable of the display and the fascia film: to do so, pull the latch out until you feel

resistance. You can then pull out the ribbon cable.

Caution!

If the latch has not been opened correctly, the ribbon cable may be damaged when it is pulled out.

- 5. Pull flow measurement hoses **56** off the flow sensor.
- 6. Pull pressure measurement hoses **57** off the pressure sensors.







- 7. Undo the four screws **51** and remove control board **50**.
- Remove spring 54 from the control board and remove battery 52 (see "7.9 Replace battery on control board" on page 42). You need the metal spring for the new control board.

If it is unnecessary to replace the display, continue from step $\ensuremath{\textbf{11.}}$

- 9. Remove display 49.
- 10. Insert a new display 49.

Caution! The display may not tilt and must be straight in the guides.

- Insert a new battery 52 in the appropriate holder of the new control board and screw on spring 54 (see "7.9 Replace battery on control board" on page 42).
- 12. Put the control board in the top part of the housing and attach it with the four screws **51**.
- 13. Reattach the pressure measurement hoses **57** to the pressure sensors.

Ensure that you do not switch the hoses: connect the connection of the flow sensor marked **Hi** to the hose connector marked **Hi** at the device outlet.

14. Attach the flow measurement hoses **56** to the flow sensor.

Ensure that you do not switch the hoses: connect the connection of the flow sensor marked **Lo** to the hose connector marked **Lo** at the device outlet.

15. To prevent hoses kinking, position flow measurement hoses 56 and pressure measurement hoses 57 as shown.







- 16. Connect the ribbon cable for the fascia film and the display to the control board again:
 - pull the latch upwards.
 - push the ribbon cable into the connector.
 - press the latch back down.
- 17. Plug the connectors of the interfaces and the rotary knob into the appropriate slots of the control board.
- 18. Refit the box and filter holder (see "7.6 Replace box and filter holder" on page 35).
- 19. Refit the power board (see "7.5 Replace power board" on page 34).
- 20. Close the device (see "7.3 Close the device" on page 32).
- Check contrast and adjust if necessary (see "4.16 Test display incl. backlighting and contrast, the LEDs and the alarm" on page 20).
- 22. **Reset the date and time,** as described in the VENTI*motion*/VENTI*logic* hospital manual.

Caution!

After changing the control board, always set the date in the following sequence: 1. YEAR, 2. MONTH, 3. DAY, otherwise you may get a faulty display.

23. Test the device (see "4. Test the device" on page 13).





- ESD protected work-station
- 3 mm Allen key
- Phillips screwdriver, size 1
- 1. Open the device (see "7.2 Open device" on page 31).
- Undo the screw 53 on the battery holder and take spring 54 and seal 55 off control board 50. Put the parts on one side.
- 3. Take battery **52** out of the holder on the control board.
- 4. Put in a fresh battery **52**.
- 5. Insert spring **54** and seal **55** on the battery holder on control board **50** and fit both parts with screw **53**.
- 6. Close the device (see "7.3 Close the device" on page 32).
- 7. **Reset the date and time** as described in the VENTI*motion*/VENTI*logic* hospital manual.

Caution! After changing the control board, always set the date in the following sequence: 1. YEAR, 2. MONTH, 3. DAY, otherwise you may get a faulty display.

 Test the device (see "4. Test the device" on page 13).

7.10 Dismantle device outlet

Tools and equipment required

- ESD protected work-station
- 3 mm Allen key
- Phillips screwdriver, size 1
- 1. Open the device (see "7.2 Open device" on page 31).
- Remove the mains supply unit (see "7.4 Replace mains supply unit" on page 33).
- 3. Remove the power board (see "7.5 Replace power board" on page 34).



- 4. Remove box and filter holder (see "7.6 Replace box and filter holder" on page 35).
- 5. Remove the control board (see "7.8 Replace control board/display" on page 39).
- 6. Pull the pressure-side hose off device outlet **90**. Dispose of the pressure-side hose (only when servicing or changing patients).
- 7. Pull flow measurement hoses off the flow sensor. Dispose of flow measurement hoses **56**.
- 8. Pull pressure measurement hoses **57** off the device outlet. Dispose of pressure measurement hoses.
- Remove device outlet 90 from the top part of the housing 36. To do so, lift the tab slightly using a screwdriver and take out the device outlet.
- 10. Remove any adhesive residues from the housing.
- 11. Place a spot of glue on the lock:
 Ø ≤ 1 mm; glue WM 14946 (Loctite 4601)
 or WM 14952 (Loctite 4031).
- 12. Fit a new device outlet **90** in the top part of the housing **36**.
- 13. Put new pressure measurement hoses **57** on the relevant connectors of device outlet **90**.
- 14. Put new flow measurement hoses **56** on the relevant connectors of device outlet **90**.
- 15. Refit the control board (see section 7.8 on page 39)
- 16. Refit the box and filter holder (see section 7.6 on page 35).
- 17. Refit the power board (see section 7.5 on page 34).
- 18. Refit the mains supply unit (see section 7.4 on page 33).
- 19. Close the device (see section 7.3 on page 32).
- 20. Test the device (see "4. Test the device" on page 13).



- ESD protected work-station
- 3 mm Allen key
- 11 mm Allen key adapter

Note

If you only want to replace the dialknob, simply pull this off the spindle of the encoder and replace it by a new one. To replace the entire encoder, proceed as described below.

- 1. Open the device (see section 7.2 on page 31).
- 2. Pull dial **83** off the spindle of encoder **82**.
- 3. Unscrew the hexagon nut on the outside of the top part of the housing **36**.
- Pull the cable of encoder 82 off control board 50.
- 5. Take the encoder out of the top part of the housing from the inside.
- 6. Insert the new encoder in the top part of the housing.
- Screw the hexagon nut onto the spindle of the encoder from the outside. Tighten up the nut.
 Ensure that the spindle of the encoder turns easily.
- 8. Plug the cable of the encoder onto the control board.
- 9. Push the dial onto the spindle of the encoder up to the stop. Check whether the dial turns easily.
- 10. Close the device (see section 7.3 on page 32).
- 11. Test the device (see "4. Test the device" on page 13).



- ESD protected work-station
- 3 mm Allen key
- Phillips screwdriver, size 1
- Knife with a smooth, flat blade
- 1. Open the device (see section 7.2 on page 31).
- 2. Remove the mains supply unit (see section 7.4 on page 33).
- 3. Remove the power board (see section 7.5 on page 34).
- 4. Remove box and filter holder (see section 7.6 on page 35).
- Remove the control board and the display (see section 7.8 on page 39). Undo fascia film 33 using a knife-blade and carefully peel it off. Use a little 70 % isopropanol to degrease this part of the housing.
- 6. Peel the protective film off the inside of the new fascia film and affix fascia film **33** to this area of the housing. Ensure that the ribbon cable is pushed cleanly through the opening in the housing without any kinks.
- 7. Carefully pull the outer protective film off the new fascia film using a fingernail.
- 8. Refit the display and the control board (see section 7.8 on page 39).
- 9. Refit the box and filter holder (see section 7.6 on page 35).
- 10. Refit the power board (see section 7.5 on page 34).
- 11. Refit the mains supply unit (see section 7.4 on page 33).
- 12. Close the device (see section 7.3 on page 32).
- 13. Test the device (see "4. Test the device" on page 13).





- ESD protected work-station
- 3 mm Allen key
- Phillips screwdriver, size 1
- Torque wrench 20 120 Ncm
- 1. Open the device (see section 7.2 on page 31).
- 2. Undo screws **87** of the humidifier socket and the ferrite core.
- 3. Pull connecting cable **70** for the humidifier out upwards.
- 4. Undo screws **86** and remove the alarm (piezo buzzer).
- 5. Remove the foam for the expansion space.
- 6. Remove the plug locators for the O_2 value.

You have now dismantled all parts. Now start assembling into the new bottom part of the housing **35**.

- 7. Insert the plug locators for the O_2 value in the new bottom part of the housing.
- 8. Put the foam for the expansion space back in.
- Refit the alarm (piezo buzzer) and fix it in position with the two screws 86. Tighten the screws with a torque of 47 Ncm.
- Put connecting cable **70** into the plug locator with the radius of the plug side facing **downwards**. Both webs of the plug must be located **within** the plug locator for this.
- 11. Fix connecting cable **70** in position with lock washer **69** and screws **87** and **88**.
- 12. Close the device (see section 7.3 on page 32).
- 13. Test the device (see "4. Test the device" on page 13).





- ESD protected work-station
- 3 mm Allen key
- Phillips screwdriver, size 1
- 1. Open the device (see section 7.2 on page 31).
- 2. Remove the mains supply unit (see section 7.4 on page 33).
- 3. Remove the power board (see section 7.5 on page 34).
- 4. Remove box and filter holder (see section 7.6 on page 35).
- 5. Remove the control board (see section 7.8 on page 39).
- 6. Remove the device outlet (see section 7.10 on page 42).
- 7. Remove the lock washers from pins 79.
- Push pins **79** out of the hinges and take handle
 78 off the top part of the housing.
- 9. Remove mains input connector **80** and cable securing clip **81**.

You have now dismantled all parts. Now start assembling into the new top part of the housing **36**.

- Take the new top part of the housing 36 in your hand and insert handle 78 in the top part of the housing.
- 11. Push pins **79** into the hinges and fit the lock washers.
- 12. Fit the device outlet (see section 7.10 on page 42).
- 13. Refit the display and the control board (see section 7.8 on page 39).
- 14. Refit the box and filter holder (see section 7.6 on page 35).
- 15. Refit the power board (see section 7.5 on page 34).
- 16. Refit mains input connector **80** and cable securing clip **81**.
- 17. Refit the mains supply unit (see section 7.4 on page 33).
- 18. Close the device (see section 7.3 on page 32).
- 19. Test the device (see "4. Test the device" on page 13).



8.1 List of spare parts for VENTImotion/VENTIlogic

Note

The item numbers in the table are identical to the numbers in the text of this service and repair guide. There are gaps in the numbering because the numbers were assigned in the operating instructions, but not all parts are spare parts.

Item no.	Name	Article no.
2	Power supply cable	WM 24177
8	Plug for pressure measurement hose	WM 24115
11	Pressure measurement hose	WM 24038
12	Hose system	WM 24130
16	Filter flap	WM 24180
31	Carrying bag	WM 24888
33	Fascia film for VENTI <i>motion</i> Fascia film for VENTI <i>logic</i>	WM 24833 WM 27033
34	Cap screw M4 x 12-8.8 ST-ZN	WM 50605
35	Housing, bottom part with nameplate	WM 24887
36	Housing, top part with fascia film, VENTI <i>motion</i> Housing, top part with fascia film, VENTI <i>logic</i>	WM 24804 WM 27028
37	Filter holder, complete	WM 24150
38	Foam, expansion space	WM 24171
39 40	Set of spare filters (sufficient for a year) Coarse dust filter Fine dust filter	WM 15682 WM 24880 WM 15026
41	VENTI <i>motion</i> : Printed circuit, power board, new Printed circuit, power board, replacement VENTI <i>logic</i> : Printed circuit, power board, new Printed circuit, power board, replacement	WM 24840 WM 24849 WM 27090 WM 27099
42	Connecting cable for humidifier, ferrite core	WM 24254
43	Piezo buzzer/alarm	WM 24109
44	Mains supply unit with insulating board	WM 24213
45	Connecting cable for power board/mains supply unit	WM 24861
46	Connecting cable for power board/control board	WM 24307
47	Cable, earth	WM 24813
48	Rechargeable battery socket with cable (connecting cable for rechargeable battery socket/power board)	WM 24507
49	LC display	WM 24828
50	Printed circuit, control board, new Printed circuit, control board, reconditioned	WM 24890 WM 24837

Item no.	Name	Article no.		
51	Screw with washer KB 25 x 12 with washer 9-3-3 (for control board)	WM 24812		
52	Battery, 3 V	WM 24946		
53	Tallow-drop screw M 3 x 6 (for spring to secure battery)	WM 53020		
54	Spring for battery	WM 24953		
55	Seal	WM 1145/32		
56	Connecting hose, silicone 3 x 7, 130 mm long	WM 24835		
57	Pressure measurement hose, silicone 2.5 x 1.5, 200 mm long	WM 24324		
58	Fan box, complete, new Fan box, complete, reconditioned	WM 24832 WM 24838		
59	Labyrinth, self-adhesive	WM 24897		
60	Motor frame, self-adhesive	WM 24898		
61	Cushioning insert, lid (fan box gasket)	WM 24868		
62	Cushioning insert, box	WM 24874		
63	Fan with bearing consisting of:	WM 24848		
64 65 66 67	 fan, complete motor bearing decoupling tube threaded pin DIN 916, M4 x 6-V2A 	VVM 24834 VVM 24862 VVM 24028 VVM 50743		
	Fan, complete, new Fan, complete, reconditioned	WM 24834 WM 24839		
68	Countersunk screw ISO 7045 M4 x 6 (for fan)	WM 51393		
69	Lock washer, humidifier cable	WM 24088		
70	Connecting cable for O_2 valve)	WM 24117		
71	Service label * - for 2007 - for 2008 - for 2009 - for 2010 - for 2011 - for 2012	VVM 0438 VVM 0498 VVM 0499 VVM 0300 VVM 0609 VVM 0610		
72	Safety test label * - for 2007 - for 2008 - for 2009 - for 2010			
73	Hose, intake side	WM 24809		
74	Hose, pressure side	WM 24852		
75	Bacteria filter, complete	WM 24148		
76	Hose system, sterilizable	WM 24120		
77	Fine filter, packed	WM 15026		
78	Handle	WM 24190		
79	Pin	WM 24210		

*When ordering, quote type, device no. and year of manufacture

Item no.	Name	Article no.
80	Mains input plug	VVM 24853
81	Cable securing clip, strain relief	WM 24317
82	Encoder	VVM 24090
83	Dial	WM 24101
84	Western socket, 6-pin	VVM 24508
85	Western socket, 4-pin	VVM 24854
86	Screw WN5412 (for piezo buzzer)	VVM 50553
87	Tallow-drop screw KB 30 x 20	WM 22597
88	Tallow-drop screw KB 30 x 6	WM 22597
89	Washer	VVM 24088
90	Device outlet, complete	VVM 24025
	VENTI <i>support,</i> complete	WM 93340
	VENTI <i>support,</i> software	WM 93350
	Converter box	WM 93360
	VENTI-O ₂ , packed	WM 24200
	Operating instructions VENTImotion DE	WM 16836
	Operating instructions VENTImotion FR; NL; IT	WM 16837
	Operating instructions VENTImotion GB; ES; PL	WM 16838
	Operating instructions VENTImotion SE; DK; NO	WM 16839
	Operating instructions VENTImotion FI; GR; RU	WM 16882
	Operating instructions VENTI <i>logic</i> DE	WM 66090
	Operating instructions VENTI <i>logic</i> FR; NL; IT	WM 66097
	Operating instructions VENTI <i>logic</i> GB	WM 66091
	Short operating instructions, VENTImotion/VENTIlogic DE	WM 16833
	Short operating instructions, VENTImotion/VENTIlogic GB	WM 16834
	Short operating instructions, VENTImotion/VENTIlogic FR	WM 16835
	Service and repair guide VENTI <i>motion/</i> VENTI <i>logic</i> DE	WM 16831
	Service and repair guide VENTI <i>motion/</i> VENTI <i>logic</i> GB	WM 16832
	Service and repair guide VENTI <i>motion/</i> VENTI <i>logic</i> FR	WM 16866
	CD-ROM hospital manual for VENTI <i>motion</i> /VENTI <i>logic</i>	WM 24805
	– hospital manual DE	WM 16915
	– hospital manual GB	WM 16916
	– hospital manual FK – hospital manual IT	WM 16917
	– hospital manual NL	WM 16990
	Operating instructions VENTI-O ₂ , DE	WM 16849
	Operating instructions VENTI- O_2 FR; NL; IT	WM 16874
	Operating instructions VENTI- O_2 GB; ES; PL	WM 16875
	Operating instructions VENTI- O_2 DK; SE; NO	WM 16876
	Operating instructions VENTI-O ₂ FI; GR; RU	WM 16889

Set for changing patients

Set

WM 15263

consisting of:

- 1 coarse dust filter
- 1 fine filter
- 2 silicone hoses, 2.5 x 1.25 SI NF, 200
- 2 silicone hoses 3/7 130
- 1 hose, pressure side
- 2 labyrinths
- 1 decoupling tube
- 1 motor frame
- 1 cushioning insert for fan box
- 1 fan box gasket
- 1 filter holder gasket
- 1 hose, intake side
- 1 foam for expansion space
- 1 foam for filter holder
- 1 device outlet, complete
- 2 countersunk screws ISO 7045-M4x6
- 1 control board, replacement
- 1 "reconditioned" label

Important!

The removed control board must be returned to Weinmann, otherwise you will be billed the full price!

Maintenance kit 5,000 hours or 2 years

Set

WM 15684

consisting of:

- 1 coarse dust filter
- 1 fine filter
- 1 battery 3 V

Maintenance kit 10,000 operating hours or 4 years

Set

consisting of:

- 1 coarse dust filter
- 1 fine filter
- 2 silicone hoses, 2.5 x 1.25 SI NF, 200
- 2 silicone hoses 3/7 130
- 1 hose, pressure side
- 2 labyrinths
- 1 decoupling tube
- 1 cushioning insert for box
- 1 fan box gasket
- 1 filter holder gasket
- 1 hose, intake side
- 1 foam for expansion space
- 1 foam for filter holder
- 1 device outlet, complete
- 2 countersunk screws ISO 7045-M4x6
- 1 motor frame
- 1 battery 3 V

WM 15679

9. Tools, test equipment and disinfectants

Below is a list of all the tools and test equipment mentioned in this service and repair guide. The tools and test equipment required in detail can be found in the respective chapter.

9.1 Tools

- Phillips screwdriver size 1
- Phillips screwdriver size 2
- Slot-head screwdriver 0.5 x 3 x 100
- 11 mm Allen key adapter

- 2 mm Allen key
- 3 mm Allen key
- Knife with smooth, flat blade for loosening fascia film
- Nail scissors or punch to mark the service label

9.2 Test equipment and fixtures

 Manual pressure gauge, accuracy ± 0.25 %
 e.g. Digima premo SR type with pressure sensor 0-50 hPa

to order from Special Instruments Henkersgasse 2; 86720 Nördlingen, Germany Postfach 1451; 86714 Nördlingen, Germany Tel.: +49 9081/220-61 or -62, Fax: +49 9081/220 63.

- Torque wrench 20 120 Ncm to order from: Hoffmann GmbH Herbert-Ludwig-Straße 4 D-28832 Achim, Germany Tel.: +49 4202/5 27-0 Fax: +49 4202/5 27-15
- PC and PC software VENTIsupport WM 93340
- Test set WM 23465 incl. 23635 pressure measurement adapter
- 115 V power source, e.g. travel adapter 230 V/115 V, 200 W
- ESD protected work-station

• Flow measurement device

Flow and pressure gauge PF-300

to order from: imtmedical ag Gewerbestrasse 8 CH-9470 Buchs, Switzerland Phone: +41 81 750 66 99 Fax: +41 81 750 66 95

www.imtmedical.com

or

Type RT 200 (Timeter)

to order from: Allied Healthcare Products Inc. 1720 Sublette Avenue St. Louis, Missouri, MO 63110 USA Phone: 001-800-444-3954 Fax: 001-314-771-5183

or

Type EKU VIP ventilator tester

to order from: EKU Elektronik GmbH Feldstraße 9a D-56291 Leiningen Phone: +49 6746-1018 Fax: +49 6746-8484 www.eku-elektronik.de

or

Flow metering pipes

Rotameter 120 l/min, class 1.6 in operating mode (1013 hPa, 20 °C) with stand, hose liners, plug-in dial and titanium floats

to order from: Yokogawa Deutschland GmbH Broichhofstr. 7-11 D-40880 Ratingen Phone: +49 2102-49 83-0 Fax: +49 2102-49 83-22 E-mail: info@de.yokogawa.com http://www.yokogawa.com/de

9.3 Disinfectants

- TERRALIN
- MIKROZID LIQUID
- GIGASEPT FF

to order from: Schülke & Mayr GmbH Robert-Koch-Str. 2 D–22851 Norderstedt, Germany Tel.: +49 40 / 52 100 - 0 Fax: +49 40 / 52 100 - 318 Internet: www.schuelkemayr.de

	VENTImotion/VENTIlogic VENTImotion/VENTIlogic w VENTIclick		
Product class as per EC directive 93/42/EEC	I	a	
Dimensions WxHxD in cm	23 x 12.5 x 34	23 x 12.5 x 45.5	
Weight	approx. 4.5 kg	approx. 4.8 kg	
Temperature range – operation – storage	+5 °C to -40 °C t	o +35 °C o +70 °C	
Air pressure range	750 – 1 (corresponds to a height	100 hPa of approx. 2500 metres)	
Max. respiration pressure at an air pressure of 600 hPa	25 hPa		
Electrical rating	115 – 230 V Tolerance -2	AC, 50–60 Hz 20 % +10 %	
Power consumption during – operation – standby	230 V 115 V 0.2 A 0.4 A 0.043 A 0.068 A	230 V 115 V 0.35 A 0.7 A 0.043 A 0.068 A	
Classification as per EN 60601-1 – type of protection against electric shock – degree of protection against electric shock	Protectic Typ	on class II De B	
Electromagnetic compatibility (EMC) as per EN 60601-1-2: – radio interference suppression – radio interference immunity	EN 5 EN 61000-3-2, EN 6100 EN61000-4-8	5011 0-3-3, EN 61000-4-2 to 6, EN61000-4-11	
Mean sound pressure level/oper- ation as per EN ISO 17510 at a distance of 1 m from the device in the patient position	approx. 35 dB (A) at 20-35 hPa approx. 31 dB (A) at 20 hPa approx. 29 dB (A) at 15 hPa approx. 27 dB (A) at 12 hPa approx. 25 dB (A) at 10 hPa approx. 23 dB (A) at 7 hPa		
Sound pressure level, alarm	approx.	62 dB (A)	
IPAP pressure range EPAP pressure range CPAP pressure range Pressure accuracy Increment	6 to 3 4 to 2 ± 0.1 (1 hPa = 1 mba	85 hPa 20 hPa 20 hPa 6 hPa hPa ar ≈ 1 cm H ₂ O)	
Minimum stable limit pressure (PLS _{min}) (min. pressure in the event of a fault) Maximum stable limit pressure (PLS _{max}) (max. pressure in the event of a fault)	≥ 0 ≤ 60	hPa) hPa	
Respiratory frequency Accuracy Increment	6 to 4. ± 0.5 1 /	5 /min /min /min	

	VENTImotion/VENTIlogic	VENTI <i>motion/VENTIlogic</i> with VENTI <i>click</i>		
I:E (Ti/T): Inspiration time Increment Accuracy	20 % to 67 % of respiratory period 1 % ±1 %			
Trigger level	can be adjusted in 6 levels, separately for inspiration and expiration; in ST mode, expiration trigger can be switched off			
Speed of pressure rise	adjustable	in 6 levels		
Speed of pressure drop	adjustable	in 6 levels		
Accuracy of volume measurement	at 23 °C	:± 15 %		
Flow at max. speed at: O hPa Tolerance	300 l/min. ±15 l/min.	275 l/min. ±15 l/min.		
Flow at max. speed with bacteria filter at: O hPa Tolerance	270 l/min. ±15 l/min.	250 l/min. ±15 l/min.		
Heating of respiratory air as per HMV	2.5 °C	depending on heating level		
Pressure constant measured to DIN EN ISO 17510 in CPAP mode	at 20 hPa: at 14 hPa: at 10 hPa: at 7 hPa:	Δp ≤ 1 hPa Δp ≤ 1 hPa Δp ≤ 1 hPa Δp ≤ 0.5 hPa		
Fine filter separation level up to 2 µm	≥ 99.7 %			
Fine filter service life	1000 hours assuming normal ambient air			
Permitted humidity in operation and storage	≤ 95 % rh (no condensate formed)			
System resistance at an air flow of 60 l/min at patient connection	VENTI <i>motion</i> with hose system WM 24130 and Silentflow WM 23600	VENTI <i>motion</i> with O ₂ hose system WM 23737, VENTI <i>click</i> WM 24365 and bacteria filter WM 24148		
	$0.19 \frac{kPa \cdot s}{l} \qquad 0.29 \frac{kPa \cdot s}{l}$			

All values determined under ATPD conditions (ambient temperature and pressure, dry).

The right to make design modifications is reserved.



10.1 Diagram of pneumatic system



10.2 Safety distances

Recommended safety distances between portable and mobile HF telecommunications equipment (e.g. mobile phones) and the therapy device

Rated output of	Safe distance as a function of transmission frequency					
HF device		in m				
in W	150 KHz - 80 MHz	80 MHz – 800 MHz	800 MHz – 2.5 GHz			
0.01	0.12	0.12	0.24			
0.1	0.37	0.37	0.74			
1	1.17	1.17	2.34			
10	3.69	3.69	7.38			
100	11.67	11.67	23.33			

Further technical data are obtainable from the manufacturer, WEINMANN, on request.

The right to make design modifications is reserved.

11. Technical amendments

Technical amendment	From device no.	Date

12. Protocols

12.1 Repair and service protocol

Device master data		Maint	enance and repairs carried out as per servi	e records	
Manufacturer: Weinmann GmbH + Co. 22525 Hamburg	Total operating hours	Comparative measurement (hPa) as per patient record	Measures/Comments	Maintenance carried VENTI <i>motion/V</i> EN service guic	l out as per VIT <i>logic</i> de
Device type:				Company	
IPAP as per patient record:hPa					
EPAP as per patient record:hPa					
Frequency f:				Date S	ignature
Initial Softstart pressure:hPa				Сотралу	
Softstart time:					
Speed of pressure rise, insp.					
Speed of pressure rise, exp.					
Humidifier stage:					() it is a set of the
Mode:				Date Date	Grante
Device no.:				Company	
Inspiration trigger:					
Expiration trigger:					
Target values:					
ΔT_{72}				Date S	ignature
T ₁₃				Company	
Owner/operator:					
				Date	ignature

12.2 Test protocol

		Test protoco	ol Ventimotion/V	/entilogic	WE			
Device:	VENTImotion VENTIlogic	WM 24810 WM 27005	Device no.:	Device	no.:	Software ve	ersion:	
1. Test equip	oment		\/ \A/AA] /					
• as per	service and re	pair instructions for	Ventimotion VV/VL16	832				
 as per 	service and re	pair instructions for	Ventimotion WM 16	832				
3. Enter dev	vice data*				Mea	sured value	OK	not Ok
4. Test for co	orrectness of te	emperature calibra	tion and calibration	status*				
5. Test of baBattery	ttery voltage ⁄ voltage is ≥ 2	2,9 V, not service ke	ey sign shown on disp	blay				
6. Test of intVENTI	erface to VEN click is working	TI<i>click</i> g correctly						
7. Check tar	get pressure v	alues at 5, 20, and	1 35 hPa					
 The at test be 	nch: target val	en target value and ue ± 0,5 hPa mask	pressure: target	Targetvalue	Test bench	Mask		
value :	± 0,5 hPa			5 hPa		hPa		
				20 hPa		hPa b D-r		
				35 NPG		nra		
8. Test of flo		nt in				//min		
 Flow is 	s 100 ± 12 l/	min				I/min		
 9. Leak test measu 	red pressure is	≥ 35 hPa				hPa		
10.Test of clo • clock i	ock function s working corre	ectly						
11.Test of int • interfa	erface to VEN ce is working c	TI-O₂ correctly						
12.Test of int • interfa	erface to VEN ce is working c	Tipower correctly						
13.Noise tes	t*							
 14.Test of all device keys a encod 	keys, encoder is working cor re detected co er is working c	r and 115 V mode rrectly with 115 V p rrectly orrectly	power supply					
15.Test of dis	play including	j backlight and cor	trast, LEDs and alar	m				
 LEDs li display contra the alc without 	ght up correctly y segments ligh st is set correctl arm generates t t distortion	y it up correctly, back ly he correct alarm so	light is working corre und sequence and co	ctly an be heard				
 16.Check of mainter 	maintenance s enance sticker c	sticker data updated and c	pplied correctly		present	yes no		
Maintenance	performed:	yes 🗌 no 🗌	Final test carried out	t:	 te			
				uu		nume		
* test points of	only feasable v	vith Weinmann test	bench					

For decades Weinmann has been developing, producing and marketing medical devices for markets around the world. In cooperation with our partners we design economic health systems for diagnosis and therapy in Sleep Medicine, Home Mechanical Ventilation, Oxygen Medicine and Emergency Medicine.



Weinmann

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